

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

Synthesis of Compounds of the Type of -Alkoxy-2-oxy-chlorophosphine-3chlorobutadienes-1.3

S/079/60/030/009/016/022/XX B001/B066

ASSOCIATION -

Institut organicheskoy khimii Akademii nauk SSSR (Institute of Organic Chemistry of the Academy of

Sciences USSR)

SUBMITTED.

August 15, 1959

Card 3/3

s,079/60/030/009/022/022/XX B001/B066

AUTHORS:

Shostakovskiy, M. F., Frilezhayava, Ye. N., Tsymbal, H., and Stolyarova, L. G.

TITLE:

Stereochemistry of Addition Reactions to the Triple Bond

PERIODICAL:

Zhurnal obshchey khimii, 1960, Vol. 30, No. 9.

pp. 3143 - 3144

There are no data available in publications on the sterecchemistry of addition reactions of any reagents to the triple bonds of diacetylene or vinyl acetylene systems, nor on the stereochemistry of TEXT: the reactions of homolytic addition of thiols to acetylenes. The authors indicate that the stereochemistry of reactions of diacetylene (I) with alkyl thiols (II) giving 1-alkyl-thiobuten-1-ines-3 (III) investigated previously by them, as well as of reactions of compounds (III) with (II) giving the 1,4-dialkyl-thiobutadienes-1,3 (IV), take place according to the scheme given here. The geometric structure (IV) follows from the sulfone structure (V) which was confirmed by means of

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Stereochemistry of Addition Reactions to the Triple Bond

5/079/60/030/009/022/022/XX B001/B066

dipole moments, ultraviolet and infrared spectra, as well as by quantitative isomerization of the cis-cis and cis-trans compounds (V) into the trans-trans compounds (V), under the action of iodine. Thus, the thiols add stereospecifically in nucleophilic reactions with the C=C bonds, both in diacetylene and vinyl acetylene systems, according to the rule of "trans-addition" (Ref. 2). Under free radical conditions, the reaction does not proceed quite stereoselectively, so that mixtures of cis-cis and trans-trans isomers (IV) are formed at low temperatures. The predominance of (IV) indicates a high specific gravity of the cis addition of the radicals. At elevated temperatures, isomerization to the trans-trans compound (IV) occurs. It was also found that the reaction of ethanethiol with ethyl-thio ethine (VI) (Ref. 3) proceeds stereospecifically both under free radical and ionic conditions, and obeys the rule of "trans-addition", since (VII-cis) is the main product yielding the sulfone (VIII-cis). Under free radical conditions, thicls may react with various acetylene compounds stereochemically in different ways. This is always due to the stability of one of the geometrical forms of the radical RSCH CX which appears

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S/079/60/030/010/002/030 B001/B075

//, /340 AUTHORS:

Shostakovskiy, M. F., Chekulayeva, I. A., and Kondrat'-

yeva, L. V.

TITLE:

Reaction of Diacetylene With Amino Alcohols and Amines. II. Synthesis of N-Alkyldiamino-1,4-butadienes-1,3 and

N, N-Dialkylamino-1-buten-1-ines-3

PERIODICAL:

Zhurnal obshchey khimii, 1960, Vol. 30, No. 10,

pp. 3179-3183

TEXT: The authors had shown in Ref. 4 that the reactions of diacetylene with aliphatic amines and β -(dialkylamino) ethanols start at room temperature and proceed exothermically without a catalyst. This paper gives a detailed description of these reactions in which the nature of the initial amine plays an essential part. Thus, primary amines react with diacetylene to form N-alkyldiamino-1,4-butadienes-1,3 (I) (80% yield) according to the following formula: $2RNH_2 + CH \equiv C - C \equiv CH \longrightarrow RNHCH = CH - CH = CHNHR$ (I) (I a): $R = n - C_4H_9$;

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Reaction of Diacetylene With Amino Alcohols S/079/60/030/010/002/030 and Amines. II. Synthesis of N-Alkyldiamino- B001/B075 1,4-butadienes-1,3 and N.N-Dialkylamino-1- buten-1-ines-3

(I b): $n-C_5H_{11}$; (I v): $iso-C_5H_{11}$. Secondary amines react under analogous conditions, forming N,N-dialkylamino-1-buter.-1-ines-3 (II) (60% yield) according to the following formula: $R_2NH + CH \equiv C-C \equiv CH \longrightarrow R_2NCH = CH-C \equiv CH$ (II) (II a): $R = C_2H_5$; (II b): $n-C_4H_9$; (II v): $n-C_5H_{11}$. The addition of the amines to compound (II) requires harder conditions. The diene structure of the reaction products was proved by spectral analysis and by diene synthesis with ethyl vinyl sulfone. The structure of the products obtained by reacting diacetylene with secondary amines was proved by hydrolysis. The synthesized products are unstable liquids which are only stable when kept in sealed ampoules between -50° and -70°. Dienes (I) and (III) absorb carbon dioxide from the air and form solid, stable carbonates. The spectral analysis carried out by B. V. Lopatin confirmed the presence of the group - C_{0-}^{0} in the dienes obtained. There are 1 table and 6 references: 3 Soviet, 2 German, and 1 US.

Reaction of Diacetylene With Amino Alcohols S/079/60/030/010/002/030 and Amines. II. Synthesis of N-Alkyldiamino-B001/B075 1,4-butadienes-1,3 and N,N-Dialkylamino-1-buten-1-ines-3

ASSOCIATION: Institut organicheskoy khimii Akademii nauk SSSR

(Institute of Organic Chemistry of the Academy of Sciences

USSR)

SUBMITTED: August 4, 1959

X

Card 3/3

5.3700 only 2209, 1273

S/079/60/030/010/003/030 B001/B075

AUTHORS:

Shostakovskiy, M. F., Kondrat'yev, Kh. I., and Gorban', A.K.

TITLE:

Investigation in the Field of Synthesis and Conversions of Oxygen-containing Organosilicon Compounds. IX. Synthesis

of Organosilicon and Naphthyl-containing Acetals

PERIODICAL:

Zhurnal obshchey khimii, 1960, Vol. 30, No. 10,

pp. 3183-3186

TEXT: In Ref. 1 the authors showed that the reaction of vinyl alkyl ether with silanols led to the formation of mixed trialkyl silyl alkyl acetals:

CH₂=CHOAlk + HOSIR₃ → CH₃CH CH₁OSIR₃

The subject of the present paper is the reaction of vinylethyl, vinyliso-propyl, and vinyl-n-butyl ethers with α -naphthyl dimethyl silanol (I). Compound (I) was obtained by hydrolyzing the acetic acid ester of α -naphthyl dimethyl silanol. Shostakovskiy and his collaborators (Ref. 1) have found that, due to an ionic mechanism, vinyl alkyl ether reacted Card 1/2

Investigation in the Field of Synthesis S/079/60/030/010/003/030 and Conversions of Oxygen-containing Organo- B001/B075 silicon Compounds. IX. Synthesis of Organosilicon and Naphthyl-containing Acetals

easily with silanols in the presence of traces of mineral acids. However, organosilicon acetals were found to be formed even without a catalyst. In this case, the yield is very high, whereas in the presence of acids (Refs. 1, 2) side reactions take place, such as hydrolysis, dimerization of the silanols, and polymerization of the initial vinyl ether. The structure of the acetals obtained was proved by hydrolyzing them with 2% sulfuric acid under the formation of α-naphthyl dimethyl silanol, acetaldehyde, and the corresponding alcohol. The a-naphthyl dimethyl silyl alkyl acetals (II) are colorless, very mobile liquids which are soluble in ordinary organic solvents but insoluble in water. There are 1 table and

ASSOCIATION:

Institut organicheskoy khimii Akademii nauk SSSR

(Institute of Organic Chemistry of the Academy of Sciences

USSR)

SUBMITTED:

December 14, 1959

Card 2/2

87539

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S/079/60/030/012/027/027 B001/B064

AUTHORS:

Shostakovskiy, M. F., Sidel'kovskaya, F. P., and

Kolodkin, F. L.

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TITLE:

Sulfides Containing Lactam Cycles

PERIODICAL: Zhurnal obshchey khimii, 1960, Vol. 30, No. 12, pp.4108-4109

TEXT: Independently of recent publications (Ref.1), the authors synthesized the following compounds by reacting the sodium salts of lactams with allyl halides: N-allyl- α -pyrrolidone (I), N-allyl- ϵ -caprolactam (II), and N-allyl- α -piperidone (III) which had hitherto not been described. Their polymerization and the copolymerization of compound (I) with methyl methacrylate and methyl acrylate were studied (Ref.2). The hitherto unknown addition reaction of the mercaptans to the compounds (I) and (II), and to N-vinyl lactams was studied. When these two compounds were heated with equimolar amounts of ethyl mercaptan (IV), n-butyl mercaptan and the ethyl ester of thioglycolic acid (V) in the presence of the dinitrile of azoisobutyric acid (0.5% of the total weight), in the ampoule at 70 - 80°C, compounds of the general formula

Card 1/2

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

s/080/60/033/04/40/045

AUTHORS:

Shostakovskiy, M.F., Sidel kovskaya, F.P., Ogibina, T.Ya.

TITLE:

A Refractometric Method for the Quantitative Determination of Q -

Pyrrolidone in a Mixture With 7 -Butyrolactone

Zhurnal prikladnoy khimii, 1960, Vol 33, Nr 4, pp 978 - 980 PERIODICAL:

 α -pyrrolidone is obtained by the interaction of \emph{J} -butyrolactone with TEXT: ammonia. In the final product there are admixtures of butyrolactone. In the literature there is no method to be found for the determination of lpha -pyrrolidone in the presence of 7 -butyrolactone. For this purpose the refractometric method is proposed. Standard mixtures of α -pyrrolidone in the presence of γ -butyrolactone were prepared and their refractive indices were measured. The data obtained are shown in a table and a graph. It is evident that the refractive index increases with the concentration of pyrrolidone. On reaching a pyrrolidone content of 35 - 40% in the butyrolactone solution the average increment of the refractive index becomes a constant value, being 5.03.10-4 on the average. The method of pyrrolidone determination has an accuracy of 11%.

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S/080/60/033/04/40/045

A Refractometric Method for the Quantitative Determination of α -Pyrrolidone in a Mixture With \ \(\int \text{-Butyrolactone} \)

There are: 2 tables, 1 graph and 5 references, 3 of which are Soviet and 2 German.

ASSOCIATION: Institut organicheskoy khimii imeni N.D. Zelinskogo AN SSSR (Institute of Organic Chemistry imeni N.D. Zelinskiy of the AS USSR)

SUBMITTED: August 13, 1959

Card 2/2

5.3700(B) /

S/020/60/132/01/40/064 B011/B126

AUTHORS:

Shostakovskiy, M. F., Gracheva, Ye. P., Kayutenko, L. A.

TITLE:

Synthesis and Conversions of Trialkylsilylethinylvinylalkyl Ethers

PERIODICAL: Doklady Akademii nauk SSSR, 1960, Vol. 132, No. 1, pp. 153-156

TEXT: The object of the authors' work is to study the interaction of ethinyl-vinylalkyl ethers with trialkylchlorosilanes. The ethers mentioned in the title were synthesized via the organomagnesium derivative, which was produced, not in tetrahydrofuran (as in Refs. 13, 14), but in sulfuric ether (2), (3). Both these reactions take place under mild conditions. The trialkylsilylethinylvinylbutyl ethers that were obtained remind one, because of their chemical properties, of the ethinylvinylalkyl ethers which contain no silicon. Both are easily hydrolized with 2% H₂SO₄. The former have also, however, some peculiarities. The hydrolysis performed to detect their structure has shown that a splitting of the Si-C bond takes place (see scheme). The butin-1-al-4 that is produced by this reaction is isomerized to tetrolaldehyde. Unlike the silicon-free ethinylvinylalkyl ethers, trialkylsilylethinylvinyl ethers are not hydrogenated via PtO₂·H₂O or via Pd precipitated on calcium sulfate. Their hydrogenation succeeds only via a mixture

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Synthesis and Conversions of Trialkylsilylethinylvinyl- S/020/60/132/01/40/064 alkyl Ethers B011/B126

of 2% Pd/CaCO₃ and 5% Pt/C. Trimethylsilylbutory-4-butadiene-1,3 was obtained by a gradual hydrogenation. The latter is condensable with maleic anhydride and forms the adduct (A), from which butylalcohol and trimethylsilanol are split off. Under the conditions of the reaction the latter gives hexamethyldisiloxane. The authors have established that the adduct is a phthalanhydride. The siliconrich ethinylvinylbutyl ether cannot be converted either by heating with butanol without a catalyst or in the presence of from 1-2% of basic or acid catalyst into trimethylsilylbutin-1-al-4-acetal. There are 1 table and 17 references, 4 of which are Soviet.

ASSOCIATION: Institut organicheskoy khimii im. N. D. Zelinskogo Akademii nauk SSSR (Institute of Organic Chemistry imeni N. D. Zelinskiy of the Academy of Sciences, USSR)

PRESENTED: January 8, 1960, by B. A. Kazanskiy, Academician

SUBMITTED: December 24, 1959

Card 2/2

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; BOGDANOVA, A.V.; USHAKOVA, T.M.; LOPATIN, B.V.

Vinyl compounds in diene synthesis. Stereospecific orientation of the diene synthesis of vinyl aryl ethers condensed with cyclopentadiene in relation to the temperature. Dokl.AN SSSR 132 no.5:1118-1121 Je '60. (MIRA 13:6)

 Institut organicheskoy khimii im. N.D.Zelinskogo Akademii nauk SSSR. Predstavleno akademikom B.A. Kamanskim. (Ethers) (Cyclopentadiene)

s/020/60/134/001/010/021 BO16/B067

5,3610 ANTHORS :

Corresponding Member of the AS USSR, Shostakovskiy. M. F.

Kolodkin, F. L. Sidel kovskaya, F. P.,

TITLE:

On the Interaction Between Lactams and Diacetylene

PERIODICAL

Doklady Akademii nauk SSSR, 1960, Vol. 134, No. 1,

pp. 102-105 TEXT: It was the purpose of the present paper a) to produce valuable unsaturated compounds with conjugate systems of multiple bonds in combination with such heteroatoms as oxygen, nitrogen, and sulfur by reacting lactams with diacetylene; b) to compare the activity of acetylene with that of discetylene in their reaction with lactams. The authors studied the addition of lactams to discetylene by the example of pyrrolidone. Compared with acetylene, this reaction takes place much more readily at 20-35°C at atmospheric pressure. Sodium salt of pyrrolidone served as catalyst. In benzene medium the process takes place much more rapidly than dioxane. The isolated crystalline main product (I) of the reaction corresponded to monopyrrolidonyl butenine. Besides, small amounts of an

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On the Interaction Between Lactams and Diacetylene S/020/60/134/001/010/021 B016/B067

isomeric compound (II) were isolated. The IR spectra (taken by T. N. Shkurina and B. V. Lopatin, collaborators at the Optical Laboratory of the authors' institute) excluded the lactim, allene-, or butadiine structures. The authors concluded from an almost complete agreement between the absorption frequencies that the structures of I and II are equal. The hydrogenation product of I is identical with N-n-butyl pyrrolidone (IV). This proves that I has the structure of 1-N-(a-pyrrolidonyl)-1-buten-3-ine. This is proved by the formation of triacetyl benzene in boiling I with 5% H2SO4. The most likely cause of the differences between I and II as to the melting temperature, solubility, and lower stability of II is probably the monotropic molecular dimorphism. By hydrolyzing I and II under less rigid conditions, the carbonyl compound formed was converted into 2,4-dinitrophenyl hydrazone (DNPH) by direct addition of 2,4-dinitrophenyl hydrazine (DNP) to the reaction mixture. In this connection, the hitherto unknown 2,4-DNPH C₁₄H₁/N₂O₂ (VI) was isolated. The authors proved that (VI) is a derivative of ?12N²(α-pyrrolidonyl)-1-buten-3-one (V) which is formed as a result of the hydration of 1-N-(α -pyrrolidonyl)-1-buten-3-ine on the triple bond. Ketone V was isolated under mild conditions also without the

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On the Interaction Between Lactams and Diacetylene S/020/60/134/001/010/021

addition of DNP. The degree of conjugation in the molecule is high. The readiness of hydration of the triple bond in N-pyrrolidonyl butenine is probably connected with the interaction between the C=O group of the lactam ring and the vinyl-acetylene chain by means of the nitrogen atom. N (a pyrrolidonyl)=1-buten-3-ine adds one thiophenol molecule in the presence of azo-iso-butyric acid dinitrile, and forms 1-N-(a-pyrrolidonyl)-4 phenyithio-1,3-butadiene (VIII). Analytically pure VIII, however, is a mixture of isomers which could not be separated by crystallization. There are ! figure, I table, and 10 references: 6 Soviet, 1 US, and 1 German.

ASSOCIATION:

Institut organicheskoy khimii im. N. D. Zelinskogo

Akademii nauk SSSR

(Institute of Organic Chemistry imeni N. D. Zelinskiy

of the Academy of Sciences, USSR)

SUBMITTED:

May 4, 1960

Cars 3/3

81:667

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2209 only

\$/020/60/135/001/020/030

B016/B067

11.2210 AUTHORS:

Shostakovskiy, M. F., Corresponding Member AS USSR,

Chekulayeva, I. A., Kondrat'yeva, L. V., and Lopatin, B.V.

的人,但是他们的人就们是一种的,你只是这些一种的人的。""你是我们的一种的人的,我们就是<mark>是我们的,我们也是我们的,我们是我们的,我们也是我们的,我们就是我们的,我</mark>

TITLE:

Structure and Some Properties of the Products of Interac-

tion Between Diacetylene and Alkyl Amines n

PERIODICAL:

Doklady Akademii nauk SSSR, 1960, Vol. 135, No. 1,pp.101-104

TEXT: In studying the reaction of diacetylene with primary and secondary alkyl amines (Ref. 1) the authors observed that the N-alkyl-diamino-1,4-butadienes-1,3 and the N,N-dialkyl-amino-1-butenines-3, respectively, are the main products. The authors succeeded in isolating the geometric isomers of N,N-diethyl-amino-1-buten-1-ine-3 (I and II) from the reaction of diacetylene with diethyl amine. The chemical transformations and the data of spectral analysis prove that I and II have cis- and trans-structures, respectively. On heating, isomer I passes over into II. The UV spectrum of II is more intensive than that of I. In the IR spectrum of I, no absorption bands were observed in the range of from 800 to 1000 cm⁻¹, in the IR spectrum of II, however, an intensive absorption band is observed at 945 cm⁻¹ which is characteristic of a trans-configuration (Table 1).

Structure and Some Properties of the Products S/020/60/135/001/020/030 of Interaction Between Diacetylene and B016/B067 Alkyl Amines

Furthermore, an intensive absorption band is observed in the spectrum of substance I at 692 cm⁻¹ which is interpreted as the CH-vibrational deformation of the isomer. In spectrum II, no corresponding band exists in this region. The IR spectra of the isomers I and II were taken on a spectrophotometer of the type UR-10. The pictures showed that the bands of the double bond are split into two components. The intensities of the components are not equal. Substance I and II may only be geometrical or place isomers: $(C_2H_5)_2$ NCH=CH-C =CH and CH₂=CN(C_2H_5)₂-C =CH. In the range 885-895 cm⁻¹ and 3075-3095 cm of the IR spectrum of both substances, no absorption bands are observed which are characteristic of a terminal double bond. This confirms the cisatrans isomerism. On the basis of the investigation of products of the addition of amines (III), alcohols (IV), and mercaptans (V) to I and II, the place isomerism seems to be excluded (see Scheme). In the reaction of n-amyl amine with the isomers I and II, 1,4-amino-substituted butadienes (III) were formed under analogous conditions which

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Structure and Some Properties of the Products S/020/60/135/001/020/030 of Interaction Between Diacetylene and B016/B067 Alkyl Amines

had the same physico-chemical constants, formed the same picrates, and also had similar IR spectra. The IR spectra of products of the addition of butyl alcohol (IV) and ethyl mercaptan (V) to I had no absorption bands corresponding to the terminal double bond. This excludes the presence of this bond in the initial isomers. Hence, the addition with the formation of I is the most essential point in the reaction of diacetylene with diethyl amine. This agrees with the results obtained by the ion reaction of the thiols with diacetylene which is stereospecific and proceeds according to the method of the "trans-addition" rule. N-butyl-diamino-1,4-butadiene-1,3 (VI) with cis-cis configuration of the substituents with respect to the double bonds is the main product resulting from the reaction of diacetylene with n-butyl amine. The structure of VI was confirmed by a diene synthesis and by data of spectral analysis. IR spectra of butadiene VI in a polar and a non-polar solvent showed that the position of the absorption bands of >C=N and >C=C bonds was only slightly influenced (Table 2). The Raman spectrum showed only one line in the region of 1600 cm⁻¹. The bond >C=N (1684 cm⁻¹ in the IR spectrum) was also present Card 3/4

Structure and Some Properties of the Products 3/020/60/135/001/020/030 of Interaction Between Diacetylene and 3016/B067 Alkyl Amines

in a second substance which was formed in a small amount in the reaction of diacetylene with n-butyl amine. This substance will be further investigated. There are 1 figure, 3 tables, and 3 references: 2 Soviet and 1 US.

ASSOCIATION:

Institut organicheskoy khimii im. N. D. Zelinskogo Akademii

nauk SSSR

(Institute of Organic Chemistry imeni N. D. Zelinskiy

of the Academy of Sciences, USSR)

SUBMITTED:

July 18, 1960

Card 4/4

ARBUZOV, A.Ye., akad.; VAVILOV, S.I., akad.; VOL'FKOVICH, S.I., akad.;

KOCHINA, P.Ya., akad.; LANDSBERG, G.S., akad.; LEYBENZON, L.S.,

akad.; PORAY.KOSHITS, A.Ye., akad.; SMIRNOV, V.I., akad.; FESENKOV,

V.G., akad.; CHERNYAYEV, V.I., akad.; KAPUSTINSKIY, A.F.; KORSHAK,

V.V.; KRAVKOV, S.V.; NIKIFOROV, P.M.; PETROV, A.D.; PREDVODITELEV,

A.S.; FRISH, S.E.; CHETAYEV, N.G.; CHMUTOV, V.K.; SHOSTAKOVSKIY, M.F.;

KUZNETSOV, I.V., red.; MIKULINSKIY, S.R., red.; MURASHOVA, N.Ya.,

tekhn.red.

[Men of Russian science; essays on prominent persons in natural science and technology: Mathematics, mechanics, astronomy, physics, chemistry] Liudi russkoi nauki; ocherki o vydaiushchikhsia deiateliakh estestvoznaniia i tekhniki: matematika, mekhanika, astronomiia, fizika, khimiia. Moskva, Gos. izd-vo fiziko-matem. lit-ry, 1961.
599 p. (MIRA 14:10)

1. Chleny-korrespondenty AN SSSR (for Kapustinskiy, Korshak, Kravkov, Nikiforov, Petrov, Predvoditelev, Frish, Chetayev, Chmutov, Shostakovskiy).

(Scientists)

SHOTAKOVSKIY, M.F.; SKVORTSOVA, C.G.; SAMOYLOVA, M.Ya.; ZAPUHHAYA, K.V.; KOSITSYNA, E.I.

Vinyl compounds. Izv.Sib.otd.All SSSk no.1:36-43 '61. (NI A 14:2)

1. Vostochno-Sibirskiy filial Sibirskogo otdeleniya All SSSR. (Vinyl compounds)

SHOSTAKOVSKIY, M.F.; POCDAFOVA, A.V.; USHAKOVA, T.H.; LOPATIF, D.V.

Vinyl compounds in the diene synthesis. Report No. 4: Comparative characteristics of the dienophilic activity of vinyl and thiovinyl others, and optical study of the adducts obtained. Izv. AN SSSR. Otd. khim. nauk no. 1:120-127 Ja 161. (MTRA 14:2)

1. Institut organicheskoy khimii im. N.D. Zelinskogo AN SSSR. (Ethers)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SIDEL'KOVSKAYA, F.P.; ZELEHSKAYA, M.G.; SHOSTAKOVSKIY, M.F.

Lactomes and lactume. Report No. 17: Dienophilic activity of E-vinyl lactumes and of the vinyl ether of N-(f-hydroxyethyl) pyrrolidone. Izv. AK SSSR. Otd. khim.nauk no. 1:128-135 Ja '51. (MITA 14:2)

1. Institut organicheskoy khimii im. N.D. Zelinskogo AK SSSR. (Lactumes) (Ether) (Pyrrolidinone)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; UVAROVA, N.I.

Synthesis and certain properties of vinyl boronyl ether. Izv. AN SSSR. Otd. khim. nauk no.2:343-348 F 161. (MIRA 14:2)

l. Dal'nevostochnyy filial Sibirskogo otdeleniya AN SSSR i Institt organicheskoy khimii im.N.D.Zelinskogo AN SSSR. (Ether)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; SIDEL'KOVSKAYA, F.P.; ZELENSKAYA, M.G.; SHKURINA, T.N., OGIBINA, T.Ya.

Lactones and lactams. Report No.18: Reaction of vinyl lactams with hydrogen chloride and alcohols. Izv.AN SSSR Otd.khim.nauk

with hydrogen chloride and alcohols. 12v. All SSSR (MTRA 14:4) no.3:482-487 Mr 161. (MTRA 14:4)

1. Institut organicheskoy khimii imeni N.D.Zelinskogo AM SSSR. (Lactams)

20941

s/062/61/000/003/008/013 B117/B208

15,8116

2203, 1372

AUTHORS:

Shostakovskiy, M. F., Khomutov, A. M., Baykova, R. I., and

Kayutenko, L. A.

TTTLE:

Studies in the field of chemical conversions of unsaturated

and high-polymer compounds. Report 17. Synthesis of

polymers and copolymers of bis-(methyl-2-buten-1-yne-3)alkyl-

silanes

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk,

no. 3, 1961, 488-491

TEXT: The authors report on the study of polymerization and copolymerization of: bis-(methyl-2-buten-1-yne-3)diethyl silane, bis-(methyl-2-buten-1-yne-3)dimethyl silane, and bis-(methyl-2-buten-1-yne-3)methyl-propyl silane. Freshly distilled monomers were used. Copolymerization was carried out continuously for 100 hr at 60° ± 1°C. Azoisobutyric acid dinitrile was used as an initiator in a quantity of 0.2 % of the total weight of the monomer. Control experiments for investigating the polymerization of initial monomers were carried out under the same conditions.

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20941 \$/062/61/000/003/008/013 B117/B208

Studies in the field of chemical ...

Bis-(methyl-2-buten-1-yne-2)diethyl silane readily polymerizes at room temperature on the air and in the presence of initiators. The polymers are transparent, hard, and three-dimensional substances. They remain unchanged when heated to 400°C. During copolymerization with methyl methacrylate, polymers of different composition are formed, according to the concentration of the initial monomers in the reaction medium. The copolymer yields were found to decrease with increasing content of bis-(methyl-2-buten-1-yne-3)diethyl silane in the reaction medium from 10 to 25 mole%. They change little later on. The number of sılane links in the copolymer increases as its concentration in the reaction medium rises. The resultant copolymers are hard, light yellow substances with high dielectric properties: $v = 10^{17} - 10^{18}$ ohmocm. Bis-(methyl-2-buten-1-yne-3) diethyl silane was used for "cross-linking" in the polymerization of methacrylic acid and styrene. For comparison, the copolymerization of methyl methacrylate with bis-(methyl-2-buten-1-yne-3)dimethyl silane and bis-(methyl-2-buten-1-yne-3)methyl-propyl silane was studied at equal molar ratios. It was found that those copolymers have the highest yields and the highest content of silane links, which contain links of bis-(methyl-2buten-1-yne-3)diethyl silane. There are 1 figure, 5 tables, and 6

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"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

20941

S/062/61/000/003/008/013 B117/B208

Studies in the field of chemical...

references: 3 Soviet-bloc and 3 non-Soviet-bloc.

Institut organicheskoy khimii im. N. D. Zelinskogo Akademii

nauk SSSR (Institute of Organic Chemistry imeni N. D. Zelinskiy, Academy of Sciences USSR) ASSOCIATION:

November 19, 1959 SUBMITTED:

Card 3/3

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

BATUYEV, M.I.; SHMONINA, L.I.; MATVEYEVA, A.D.; SHOSTAKOVSKIY, M.F.

Optical study of the structure of some 1, 2, 3-substituted 1, 3-butadienes. Izv.AN SSSR Otd.khim.nauk no.3:513-516 Mr !61. (MIRA 14:4)

1. Institut goryuchikh iskopayemykh AN SSSR i Institut organicheskoy khimii imeni N.D.Zelinskogo AN SSSR.
(Butadiene)

S/062/61/000/003/009/013 B117/B208

15.8103

2203, 1374

Shostakovskiy, M. F. and Shapiro, E. S.

TITLE:

AUTHORS:

Synthesis of phthalyl glycine vinyl ester and some of its

conversions

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh

nauk, no. 3, 1961, 517-519

TEXT: In this news in brief, the authors report on their method of direct vinylation of N-substituted A-amino acid. Phthalyl glycine was used as standard substance. Vinylation was carried out in a rotating autoclave with a manometer and a thermocouple with automatic temperature regulation. The reaction took place at a high acetylene excess in absolute dioxane medium, i.e., under conditions that had been repeatedly applied with success by the authors. Anhydrous zinc acetate was used as catalyst. The corresponding vinyl ester was thus obtained in 80-85 % yield:

 $c_{6}H_{4} < c_{0}$ NCH₂COOCH = CH₂ is a crystalline substance insoluble in most

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20942 \$/062/61/000/003/009/013 B117/B208

Synthesis of phthalyl glycine vinyl

organic solvents, in petroleum, and in water. This fact is used for purifying the vinyl ester from the initial phthalyl glycine in those cases in which vinylation does not quantitatively proceed. Some conversions of the respective vinyl ester were studied. Its hydrolysis was found to occur under more rigorous conditions than in vinyl alkyl esters, i.e., on heating to 80°C with 15 % sulfuric acid. Thiols are added smoothly and quantitatively during catalysis with a radical initiator, thus giving the corresponding sulfides: phthalyl glycine-8-ethyl-thioethyl ester $C_{14}H_{15}O_4NS$, boiling point 1390-1410C (0.015 mm); phthalyl glycine- β -butylthioethyl ester $^{\rm C}_{16}{}^{\rm H}_{19}{}^{\rm O}_4{}^{\rm NS}$, boiling point 150°-152°C (0.008 mm). After distillation, both sulfides become nearly colorless, viscous liquids which crystallize on standing. After recrystallization from petroleum ether, their melting points are 43° and 37°-38°C, respectively. In the presence of the dinitrile of azoisobutyric acid and benzoyl peroxide, the phthalyl glycine vinyl ester is polymerized. Also its copolymerization with methyl methacrylate was shown to be possible in principle. There are 3 tables and 6 references: 2 Soviet-bloc and 4 non-Soviet-bloc.

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"APPROVED FOR RELEASE: 08/09/2001

CIA-RDP86-00513R001549920002-9

2091.2

s/062/61/000/003/009/013 B117/B208

Synthesis of phthalyl glycine vinyl ...

Institut organicheskoy khimii im. N. D. Zelinskogo Akademii ASSOCIATION:

nauk SSSR (Institute of Organic Chemistry imeni N. D. Zelinskiy,

Academy of Sciences USSR)

SUBMITTED:

May 4, 1960 (initially) October 9, 1960 (after revision)

Card 3/3

CIA-RDP86-00513R001549920002-9" APPROVED FOR RELEASE: 08/09/2001

CHEKULAYEVA, I.A.; SHOSTAKOVSKIY, M.F.; GLADYSHEVSKAYA, V.A.; LIPOVICH, I.V.

Synthesis and transformations of vinyl ethanolamine ethers. Part 13: Copolymerization of some vinyl ethanolamine ethers with methacrylate. Vysokom.soed. 3 no.6:901-907 Je 161. (MIRA 14:6)

1. Institut organicheskoy khimii imeni N.D.Zelinskogo. (Ethanol) (Methacrylic acid) (Polymerization)

SHOSTAKOVSKIY, M.F.; LASKORIN, B.N.; NIKUL'SKAYA, G.N.; CHEKULAYEVA, I.A.; IOANISIANI, P.G.

Suspension polymerization of the trivinyl ether of triethanolamine.

Synthesis of a new anion exchanger. Vysokom.soed. 3 no.6:908-911

Je 161.

1. Institut organicheskoy khimii imeni N.D.Zelinskogo AN SSSR. (Ethanol) (Ethers) (Polymerization) (Ion exchanger resins)

s/190/61/003/007/004/021 25260 B101/B208 15.8050 Shostakovskiy, M. F., Sidel'kovskaya, F. P., Ibragimov, F. AUTHORS: Copolymerization of vinyl pyrrolidone and vinyl caprolactam TITLE: with dimethyl vinyl ethinyl carbinol Vysokomolekulyarnyye soyedineniya, v. 3, no. 7, 1961, PERIODICAL: 976-979 TEXT: The purpose of the present paper was to study the fundamental. rules governing the copolymerization of vinyl pyrrolidone (VP) and vinyl caprolactam (VC) with dimethyl vinyl ethinyl carbinol (CARB). It was of interest in this connection that CARB is the raw material for the so-called carbinol glues. The following formula is given for the structure of the copolymers: ... - CH₂ - CH - ...C (CH₃)₂ OH n = 3; 5.Card 1/5

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

		2526G	" s/190/61/	003/007/004/0	21
o-malumomi	ration of wire	pyrrolidone	. B101/B208	3	
For the li	aks of the con	oolymer which con	sist of carbino	ol, also the	40
					trile 1
					1
was added	as initiator.	The results for	transport	tion of the	1
initial mi	vture, mole-%	yield of copoly	mer, composition	er, mole-%	
•			VC VC	CARB	.15
VC	CARB				
.100	0	76.5	100.0		
90	10	18.7	65.7	34.3	
75	25 '	19.7	38.1	61.9	•
75 50	50 ·	33.6	12.6		50
25	75	59.1	. 4.8		
10	90 ·	60.7	was not	determined 100.0	
0	100	97•5	. 0		
	ition of the	copolymer was cal	culated from i	ts nitrogen c	ontent.
Mb	TATOM OF AMO	for VP + CARB:			5
The compos	ing was found				* * * * * * * * * * * * * * * * * * * *
The compose The follow	ing was found		*	,	
The compose The follow	ing was found		; ·		
The compose The follow Card 2/5	ring was found		•		

Copolymerization of vinyl pyrrolidone ... S/190/61/003/007/004/021 B101/P208 ... composition of the

initial		yield of copolymer, %	copolymer, cole-		
100 90 75 50 25	0 10 10 25 50 75 90	67.5 13.7 23.4 27.9 61.3 76.5	100.0 46.2 27.9 9.3 0.5 was not	53.8 .72.2 .0.0 .75.5 detormined .100.0	

The following conclusions say be drawn: 1) The copolymers constained were CARB than the initial mixture; 2) the yield increased with increasing CARB content. The copolymers of both types were soluble in scotone and others, insoluble in diethyl ether, petroleum ether, chlorofort, benzine and water. Carticularly noted is the poor solubility in meating of the copolymer from 10,1 ve and 90 100 MB. Studying the solubility of the copolymers and homopolymers in some nitrogen-containing solvente:

Card 3/5

		3/190/61/003/	007 4 / 621	
· Copolymerization of vir	25260 nyl pyrrolidone	3101/B208		
95 1 VO - 75,3 0.498 10 - VO - 903 03 U	olymers: 10 the lymer. The rol mer solutions: relative viscosia/, 100 5.30d 1.111 1.532 1.921	composition of the composition of the composition of the nittal mixture nomopolymer V: 500 VP 500 CARB 250 VP 750 OARB 100 VP 260 CARB	with the 70 or e relative relative vicability,2 0.513	r c
The copolymers have ad with increasing CaRb of Soviet-bloc references	ontont. There	n-forming properties of are 2 figures, 5 table	nich increase es, ena 2	
Card 4/5				

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

25260 S/190/61/003/007/004/021 B101/B208

Copolymerization of vinyl pyrrolidone ...

,

ASSOCIATION: Institut organicheskoy khimii im. N.D. Zelinskogo AN SSSR

(Institute of Organic Chemistry imeni N.D. Zelinskiy,

AS USSR)

SUBMITTED:

August 7, 1960

Card 5/5

26290 \$/190/61/003/008/003/019 B110/B220

15.8150

AUTHORS: Shostakovskiy, M. F., Kotrelev, V. N., Kalinina, S. P.,

Kuznetsova, G. I., Layne, L. V., Borisova, A. I.

TITLE:

Organotin monomers and polymers. IV. Synthesis and conversion

of tin-containing esters of acrylic and cinnamic acids

PERIODICAL:

Vysokomolekulyarnyye soyedineniya, v. 3, no. 8, 1961,

1128-1130

TEXT: The present paper deals with the synthesis of organotin derivatives of cinnamic and acrylic acids. The synthesis was performed by a method developed by the authors. The vaporous alkyl halide was reacted in a tube furnace or autoclave with an Sn-Mg alloy in the presence of various solvents and catalysts. The alkyl-halide tin compounds formed were saponified with lye to the corresponding hydroxy derivatives, and then the esters were obtained by reaction with acrylic or cinnamic acid. 1) Triethyl-stannyl acrylate $(C_2H_5)_3$ SnOCOCH=CH2, was obtained from a 50% aqueous solution of acrylic acid at 5-10°C by adding triethyl stannol. The white crystalline

Card 1/3

Organotin monomers and polymers ...

26290 S/190/61/003/008/003/019

precipitate (melting point 102°C) could be dissolved in organic solvents. 2) In the same way, tributyl-stannyl acrylate was obtained from hexabutyl stannous oxide and acrylic acid. 3) The triethyl-stannyl ester of cinnamic acid was obtained from cinnamic acid and hexaethyl stannous oxide according to the equation $(c_2H_5)_6Sn_2O + 2C_6H_5 = CHCOOH \longrightarrow 2(c_2H_5)_3SnococH = CHC_6H_5$ + H20. The organotin compounds obtained polymerize easily, and form transparent solid copolymers with styrene and methyl methacrylate. The thermomechanical properties of some polymers and copolymers are shown in Fig. 2.

ASSOCIATION: Nauchno-issledovatel'skiy institut plasticheskikh mass (Scientific Research Institute of Plastics)

SUBMITTED:

September 1, 1960

Card 2/3

761/003/008/004/019 B110/B220

15.6150

Shostakovskiy, M. F., Kotrelev, V. N., Kuznetsova, G. I.,

Kalinina, S. P., Layne, L. V., Borisova, A. I.

AUTHORS:

Studies on the synthesis and conversions of organotin monomers and polymers. V. Study of the formation of organotin polymers as a function of the polymerization conditions, and some physicochemical properties of organotin TITLE:

Vysokomolekulyarnyye soyedineniya, v. 3, no. 8, 1961, polymers

PERIODI CAL:

TEXT: The present study deals with the yield in polymers of triethyl-TEAT: The present study dears with the yierd in polymers of threthyrs stannyl methacrylate and acrylate as a function of polymerization time, svannyl methacrylate and acrylate as a lunction of polymerization time, temperature, initiation, and concentration. Benzoyl peroxide, azoisobutyric demperature, initiation, and concentration. Benzoyi peroxide, azoisobutyri acid dinitrile, or triethyl-benzyl ammonium chloride served as initiators. acid dinitrile, or triethyl-penzyl ammonium chioride served as initiator;
The results are shown in Fig. 1. The composition of the copolymer from
triethyl-stannyl methacrylate and methyl methacrylate was studied for
initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.10 At an initial molar ratios of the components of 1.4 4.4 and 4.4 triethyr-standyr methacryrate and methyr methacryrate was studied for initial molar ratios of the components of 1:1, 1:4, and 1:12. At an initial

Card 1/5

s/062/61/000/004/006/008 B118/B208

5.3700 2209

Shostakovskiy, M. F., Komarov, N. V., Kuznetsova, V. P., and

Study in the field of synthesis and conversions of unsaturated AUTHORS: TITLE

organosilicon compounds. 1. Esterification of primary and tertiary 7-silicon-containing acetylene alcohols by adipic

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk, PERIODICAL:

no. 4, 1961, 699-703

TEXT: The authors studied the esterification of primary and tertiary 7-silicon-containing acetylene alcohols with adipic acid. This esterification was accomplished by direct reaction of the alcohols with adipic acid, and by reaction of these alcohols, or their magnesium derivatives, with adipic acid chloride. The esterification of the primary 7-silicon-containing acetylene alcohols with adipic acid without catalysts takes place R_3 SiC=C-CH₂OH + HOOC(CH₂)₄COOH $\rightarrow R_3$ SiC=C-CH₂OOC(CH₂)₄COOH, where

Card 1/4

Study in the field ...

S/062/61/000/004/006/008 B118/B208

 $R = CH_3$, C_2H_5 , and C_6H_5 , or to form the neutral adipic ester; $2R_3SiC = C-CH_2OH + HOOC(CH_2)_4COOH \rightarrow R_3SiC = C-CH_2OOC(CH_2)_4COOCH_2C = CSiR_3$, where $R = CH_3$ and C_6H_5 , depending on the conditions and the quantitative ratio of the components. This method is, however, not applicable to the esterification of tertiary γ -silicon-containing acetylene alcohols, since the reaction of these alcohols with adipic acid, with or without acid catalysts (such as boric acid etc.) gives rise to dehydration of the acetylene hydrocarbons:

 R_3 SiC=C-C-OH---> R_3 SiC=C-C-CH₂ + H_2 O, where R = CH_3 and C_2H_5 . Also

the esterification of tertiary 7-silicon-containing acetylene alcohols by their reaction with adipic acid chloride in the presence of pyridine as under the action of sodium ethylate were unsuccessful. The synthesis of Card 2/4

X

Study in the field...

S/062/61/000/004/006/008 B118/B208

acid and neutral esters of tertiary y-silicon-containing acetylene alcohols was accomplished by reaction of magnesium alcoholates with adipic acid chloride

. СН_а

 $R_3SiC \equiv C - COM_gBr + CIOC(CH_2)_1COCI \rightarrow$

ĊH_p .

 CH_3 $\rightarrow R_3SiC \equiv C - COOC(CH_2)_1COCI \xrightarrow{\div H_3O} R_3SiC \equiv C$

—соос (СН₂),соон

 CH_3

Card 3/4

Study in the field...

$$CH_3$$
 $2R_3SiC = C - COM_gBr + CIOC(CH_2)_4COCI \rightarrow CH_3$

22517 5/062/61/000/004/006/008 B118/B208

$$CH_3 \qquad CH_3$$

$$\rightarrow R_3SiC \equiv C - COOC(CH_2)_1COOC - C \equiv CSiR_3,$$

$$CH_3 \qquad CH_3$$

$$CH_3 \qquad CH_3$$

$$CH_3 \qquad CH_3$$

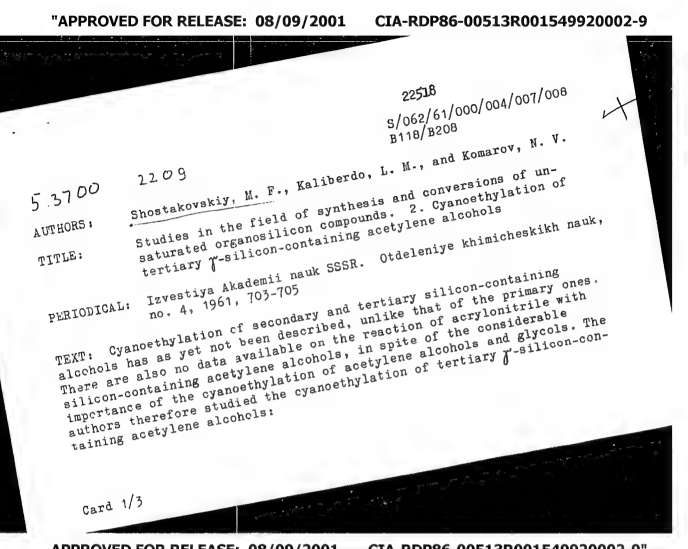
There are 7 Soviet-bloc references.

ASSOCIATION: Irkutskiy institut organicheskoy Ehimii Sibirskogo otdeleniya AN SSSR (Irkutsk Institute of Organic Chemistry

of the Siberian Branch of the Academy of Sciences USSR)

SUBMITTED: December 15, 1959

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CIA-RDP86-00513R001549920002-9" APPROVED FOR RELEASE: 08/09/2001

Studies in the field ...

S/062/61/000/004/007/008 B118/B208

$$R_{3}SiC = C - CH_{3} - CH_{2} - CH_{2} - CH_{3} - CH_{3}SiC = C - CH_{2}CH_{2}CN, \text{ where } R = CH_{3} \text{ and } CH_{3}$$

 c_{2}^{H} . This cyanoethylation of acetylene alcohols was carried out under the action of a 40% aqueous caustic potash solution and also of sodium methylate. The catalytic effect of caustic potash solution gave no β -cyanoethyl ethers. The application of sodium methylate permitted the synthesis of β -cyanoethyl ethers of the silicon-containing acetylene alcohols used (50-60% yields). The reaction is exothermic. Better yields are obtained by using benzene as the solvent. The reaction of the β -cyanoethyl ether of 4-trimethyl-silyl-2-methyl butyn-3-ol-2 with ethyl magnesium bromide gave. like several analogous cases, the corresponding silicon acetylene ketone according to the equation

$$\begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{CH}_{3} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \\ \text{C} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{SiC} = \text{C} - \begin{array}{c} \text{CH}_{3} \\ \text{C} \end{array})_{3} \text{C} \times \text{C}$$

Card 2/3

Studies in the field...

S/062/61/000/004/007/008 B118/B208

presence of a carbonyl group in this compound was confirmed by the formation of 2,4-dinitro-phenyl hydrazone. In addition to the formation of the erganosilicon ketone according to this equation also a cleavage of the β -cyanoethyl ether takes place to give 4-trimethyl-silyl-2-methyl-butyn-3-ol-2. Other conversions of the β -cyanoethyl ethers of tertiary resilicon-containing acetylene alcohols will be described in the following report. There are 8 references: 4 Soviet-bloc and 4 non-Soviet-bloc. The two references to English-language publications read as follows: R. Cooksen, F. Mann, J. Chem. Soc. 618 (1947), 67 (1949); S. Nozakura, S. Konotsune, Bull. Chim. Soc. Japan 29, 322, 326 (1956).

ASSOCIATION: Institut khimii Vostochno-Sibirskogo filiala Sibirskogo otdeleniya AN SSSR (Institute of Chemistry of the Eastern Siberian Branch of the Siberian Department, Academy of

SUBMITTED: December 15, 1959

Card 3/3

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOTHINGT REP. CHERRIOR, L.M.; ALLION, L.P.

Chemical conversions of unsaturated and high molecular weight compounds. Report No. 18: Polymerization and copolymerization of divinyl tartrate and methyl methacrylate. Izv.AN SSSR Otd.khim.nauk no.4: 706-709 Ap 161. (MIRA 14:4)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Tartaric acid) (Methacrylic acid)

SHOSTAKOVSKIY, M.F.; BOGDANOVA, A.V.; USHAKOVA, T.M.

Diene synthesis with the participation of vinyl ethers. Report No. 5: Reaction of divinyl ether with hexachlorocyclopentadiene. Izv.AN SSSR Otd.khim.nauk no.4:709-714 Ap *61. (MIRA 14:4)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Ether) (Cyclopentadienė)

SHOSTAKOVSKIY, M.F.; BOGDANOVA, A.V.; PLOTNIKOVA, G.I.

Study of diacetylene derivatives. Report 10.8: Reactions of ethydyl vinyl and thioethynyl vinyl ethers with carbonyl compounds. Izv. AN SSSR. Otd. khim.nauk no.5:905-909 My 161. (MIRA 14:5)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Ether) (Carbonyl compounds)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; SIDEL'KOVSKAYA, F.P.; ZELENSKAYA, M.G.

Lactones and lactams. Reprot No.19: Synthesis of ethers and esters of N-(A-hydroxyethyl)pyrrolidinone. Izv.AN SSSR.Otd.khim.nauk no.5:

(MIRA 14:5)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Pyrrolidinone)

23591 \$/062/61/000/005/008/009 B118/B220

15.8102 2209

AUTHORS; Shostakovskiy, M. F., Gracheva, Ye. P., Meshcheryakov, A. P., and Glukhovtsev, V. G.

TITLE:

Polymerization of the vinyl ether of methyl cyclopropyl carbinol

PERIODICAL: Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh nauk, no. 5, 1961, 924 - 927

TEXT: In Ref. 1 (B. A. Zakharov et al., Dokl. ANSSSR, 122, no. 5, 814 (1958)), it has been stated that the double bond of the vinyl ethers has an increased nucleophilic character which manifests itself in various addition reactions, transformations, and especially in the polymerization reaction. For the study of the conditions of polymerization of the compounds CH₂ = CHOR (I), the vinyl ether of methyl cyclopropyl carbinol is

 $CH_2 = CH-O-CH-CH \int_{CH_3}^{CH_2} (II).$

According to the rule of Markovnikov, the cyclopropyl group of this ether,

23591 \$/062/61/000/005/008/009 B118/B220

Polymerization of the ...

as possible carrier of the propenyl group, is able to add various polar compounds. Moreover, this ether may be of interest as test substance for the synthesis of different polymers in the polymerization and copolymerization reactions. The present paper describes the polymerization of the vinyl ether of methyl cyclopropyl carbinol in the presence of the initiators FeCl₃ and azonitrile isobutyric acid under optimum conditions for

the polymerization of the vinyl alkyl ethers. It has been found that compound (II) shows higher reactivity during polymerization in the presence of a 5 % solution of iron perchloride (in dioxane) than vinyl alkyl ethers (I) under the same conditions. First of all, this is evident from the fact that the polymerization of the ether (II) begins at 0°C and the highest yield in polymer is obtained at a temperature of -17 to -20°C whereas other vinyl alkyl ethers polymerize at boiling temperature only. The reason for such diverging temperatures of polymerization is the different stability of the ozonium complexes of these compounds (I, II):

(2017) My - May 1

H(FeC1_OH{

OH = CH-Q-CH-CH

 $\langle z c c \rangle$

Card 2/3

"APPROVED FOR RELEASE: 08/09/2001

CIA-RDP86-00513R001549920002-9

Polymerization of the ...

Evidently, complex (IV) is of lower stability; its decomposition is effected at a low temperature resulting also in the formation of a polymer at lower temperature. The use of azonitrile isobutyric acid as initiator instead of FeCl₃ did not give any results. There are 3 Soviet-bloc references.

ASSOCIATION: Institut organicheskoy khimii im. N. D. Zelinskogo Akademii nauk SSSR (Institute of Organic Chemistry im. N. D. Zelinskiy, Academy of Sciences USSR)

SUBMITTED: October 12, 1960

Card 3/3

25039 \$/062/61/000/006/001/010 B118/B220

5.3400

Shorygin, P. P., Shkurina, T. N., Shostakovskiy, M. F., and

Gracheva, Ye. P.

TITLE:

AUTHORS:

Spectra and structure of vinyl ethers

PERIODICAL:

Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh

nauk, no. 6, 1961, 1011 - 1015

TEXT: A study has been made of the influence of various functional groups on the properties of vinyl compounds and on the state of the double bond C=C. In the case of vinyl ethers, the influence exerted by the alkoxy group upon the double bond becomes evident in chemical properties, such as increased reactivity in addition reactions, in hydrolysis, and in polymerization and copolymerization processes. Not much material is available in the literature with regard to the influence of the alkoxy group upon the strength of the double bond C=C and upon the optical properties. The present paper deals with studies concerning the Raman spectra and the ultraviolet absorption spectra in vacuo for a series of vinyl ethers containing alkyl, naphthene and aromatic radicals. Most vinyl ethers

Card 1/6

25039 S/062/61/000/006/001/010 B118/B220

Spectra and structure of ...

Bull. Chem. Soc. Japan 29, 110 (1956).

have several lines in the frequency range of the stretching vibrations C=C. The splitting depends on the branchings of the alkyl group and on the presence of an &-substituent. The influence of temperature upon the intensity of the lines \sim 1610 and \sim 1640 cm⁻¹ of vinyl butyl ether indicates their relation to the two rotational isomers. The influence of the group OAlk on $\omega_{\text{C=C}}$ may be regarded qualitatively as a tendency to reduce the frequen-The wavelength of the first absorption band and the intensity of the C=C line in the Raman spectra are greater for vinyl ethers than for alkenes of similar structure. The influence of solutions and temperature on the structure of the C=C band was studied. Thus, the geometric configuration of the molecules of vinyl ethers may be important to both the physical and chemical properties. The Raman spectra were taken with the NCT-67 (ISP-67) spectrograph with the Hg line 4358A. The absorption spectra were taken with spectrophotometers of types $C\Phi$ -4 (SF-4) and $C\Pi$ -41 (SP-41) with the assistance of V. A. Petukhov. There are 2 figures, 1 table, and 3 references: 1 Soviet-bloc and 2 non-Soviet-bloc. The 1 reference to English-language publication reads as follows: Y. Mikawa,

Card 2/6

"APPROVED FOR RELEASE: 08/09/2001

CIA-RDP86-00513R001549920002-9

25039 S/062/61/000/006/001/010 B118/B220

Spectra and structure of ...

Institut organicheskoy khimii im. N. D. Zelinskogo Akademii

nauk SSSR

(Institute of Organic Chemistry imeni N. D. Zelinskiy of the

Academy of Sciences USSR)

SUBMITTED:

ASSOCIATION:

June 1, 1959

Fig. 1: Ultraviolet absorption spectra of octene-1 and vinyl ethers (solutions in heptane). Legend: bu-butyl group; >-cyclohexyl group.

Card 3/6

CIA-RDP86-00513R001549920002-9" APPROVED FOR RELEASE: 08/09/2001

SHOSTAKOVSKIY, M.F.; SIDEL'KOVSKAYA, F.P.; ROGOVA, E.V.; KOLODKIN, F.L.;

Lactones and lactams. Report 20; Reactions of N-(chloralkyl) lactams with alcohols. Izv.AN SSSR.Otd.khim.nauk no.6:1111-1116
Je '61. (MIRA 14:6)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Lactams) (Alcohols)

SHOSTAKOVSKIY, M.F.; KUZNETSOV, N.V.; DUBOVIK, N.A.; ZIKHERMAN, K.Kh.

Synthesis of ethoxyacetalaldehyde and its chemical transformations.

Izv. AN SSSR. Otd.khim.nauk no.8:1495-1500 Ag '61.

(MIRA 14:8)

1. Irkutskiy institut organicheskoy khimii Sibirskogo otdeleniya
AN SSSR.

(Acetaldehyde)

261,05 s/062/61/000/008/010/010 B117/B206

15.805D

Card 1/4

AUTHORS:

Shostakovskiy, M. F., Sidel'kovskaya, F. P., Shapiro, E. S.,

and Ogibina, T. Ya.

 β -(N-pyrrolidonyl) ethylvinyl sulfide

Akademiya nauk SSSR. Izvestiya. Ctdeleniye khimicheskikh TITLE:

nauk, no. 8, 1961, 1524-1526 PERIODICAL:

TEXT: The authors investigated the vinylation of the previously prepared N-(β-mercaptoethyl) pyrrolidone (Ref. 1: M. F. Shostakovskiy, F. P. Sidel'kovskaya, E. S. Shapiro, T. Ya. Ogibina, Izv. AN SSSR. Otd. khim. n., 1958, 68). The reaction was carried out in dioxane medium with a 2- to 4-fold acetylene excess. A rotating autoclave (250 ml) fitted with manometer, thermocouple, and automatic temperature control was used. Vinylation proceeds smoothly and with good yield (81.8 %) in the presence of 10 % caustic potash. β -(N-pyrrolidonyl) ethylvinyl sulfide (I) is a colorless, weakly smelling, viscous liquid, practically soluble in any organic solvent. Some of its conversions were investigated: addition of thiols, polymerization, and copolymerization. The addition

26405 \$/062/61/000/008/010/010 B117/B206

 β -(N-pyrrolidonyl) ethylvinyl sulfide

of the thiols is practicable during radical initiation (azoiscoutyric acid dinitrile). Corresponding sulfides are formed thereby with good yield. Addition of ethyl-thiol produces 88 % β-pyrrolidenyl ethyl-βethyl mercapto ethyl sulfide with boiling point 117-120°C (0.0:5 mm); 1.5440; d_A^{20} 1.1222. During heating the synthetized monomer (I) undergoes thermal polymerization. This is accelerated by addition of azoisobutyric acid dinitrile. The new polymer is a transparent, almost colorless, semisolid product. It is soluble in water, alsohol, benzene, and other common organic solvents with the exception of disthyl- and petroleum ether. The monomer (I) does not only form homopolymers, but participates also in the copolymerization with other vinyl monomers. was found to be extremely active. According to its activity, it is similar to acrylonitrile and methyl acrylate. It is of much higher reactivity than vinyl acetate and vinyl pyrrolidone. Polymerization and copolymerization occurred under standard conditions: in ampullas at 60°C within 100 hr in the presence of azoiso butyric acid dinitrile. Diethyl ether was used for the precipitation of polymers and copolymers. Petroleum ether was only used for copolymers of (I) and methyl acrylate.

Card 2/4

26h05 S/062/61/000/008/010/010 B117/B206

 β -(N-pyrrolidonyl) ethylvinyl sulfide

The results are listed in the Table. There are 1 table and 5 references: 4 Soviet and 1 non-Soviet.

ASSOCIATION: Institut organicheskoy khimii im. N. D. Zelinskogo Akademii nauk SSSR (Institute of Organic Chemistry imeni

N. D. Zelinskiy, AS USSR)

SUBMITTED: February 22, 1961

Table: Properties of the polymers produced. Legend: 1) Designation;
2) appearance; 3) yield, %; 4) determined S, %; 5) content of (I) links in
2) appearance; 3) yield, %; 4) determined S, %; 5) content of (I) links in
3) the copolymer, mole%; 6) solubility; 7) acetone; 8) dimethyl formamide;
4) sulfuric ether; 10) petroleum ether; 11) homopolymer of vinyl sulfide
4) sulfuric ether; 10) petroleum ether; 11) homopolymer of (I) and
4) copolymer of methylacrylate and (I); 13) copolymer of (I) and
5) copolymer of (I) and vinyl pyrrolydone; 15) copolymer
7) vinyl acetate; 14) copolymer of (I) and vinyl pyrrolydone; 15) copolymer
8) vinyl acetate; 14) copolymer of (I) and vinyl pyrrolydone; 15) transparent,
8) of (I) and acrylonitrile; 16) transparent, elastic mass; 17) transparent,
8) semisolid product; 18) transparent, elastic product; 19) white, hard,
8) semisolid product; 18) transparent, elastic product; 19) white, hard,
8) brittle. *) for C8H13ONS 18.72 % S were calculated. **) P = soluble;
8) H = insoluble; P.OPP. = restrictedly soluble.
8) Card 3/4

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHORYGIN, P.P.; SHOSTAKOVSKIY, M.F.; PRILEZHAYEVA, Ye.N.; SHKURINA, T.N.; STOLYAROVA, L.G.; GENICH, A.P.

Structure and spectra of vinyl sulfides. Izv. AN SSSR. Otd.khim.nauk (MIRA 14:9)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Vinyl sulfide-Spectra)

S/062/61/000/009/011/014 B117/B101

5.3400

AUTHORS:

Shostakovskiy, M. F., Kuznetsov, N. V., and Yang Che-Min

TITLE:

Synthesis and transformations of new vinyl ether derivatives

PERIODICAL:

Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh

nauk, no. 9, 1961, 1685-1688

TEXT: The present work was undertaken as a part of the studies of synthetic methods for the preparation of new vinyl ether derivatives. In the course of it, several new 3-hydroxy-acrolein ethers and esters were prepared by the following reactions.

Card 1/5

27**៤95** \$/062/61/000/009/011/014 B117/B101

Synthesis and transformations ...

$$CH (OC_2H_6)_3 + CH_2 = CHOC_2H_6 \xrightarrow{BF_3 \cdot (C_1H_1)_3O} (C_2H_6O)_2 CHCH_2CH (OC_2H_6)_2$$

$$CH_3OCH = CBrCHO \xrightarrow{CH_1N_2} HOCH = CBrCHO$$

$$(II) \qquad \qquad (II)$$

$$CH_3COOCH = CBrCHO \xrightarrow{CH_3COC1} N_3OCH = CBrCHO \rightarrow (IV) \qquad (III)$$

$$C_4H_5COC1 \xrightarrow{C_4H_5COC1} C_6H_6COOCH = CBrCHO \rightarrow (V)$$

The authors investigated whether these 3-hydroxy acrolein ethers and esters give the diene condensation. The ether (II) and esters (IV) and (V) obtained readily hydrolyze under the influence of water and atmospheric moisture. They consolidate on storage. Diene condensations of (II), (IV), and (V) with vinyl-butyl ether could not be realized, since the reaction products resinified completely, probably as a result of hydrogen bromide formation. Ethyl orthoformiate, however, reacts readily with dioxene in Card 2/5

27495 \$/062/61/000/009/011/014 B117/B101

Synthesis and transformations ...

the presence of boron trifluoride etherate, yielding 2-ethoxy-3-diethoxy-methyl -p-dioxane (VI) ($C_{11}H_{22}O_5$, b.p. 135°-137°C (20 mm Hg), n_D^{20} 1.4337). This may be hydrolyzed to 2-formyl p-dioxene (VII) ($C_5H_6O_3$, m.p. 40° - 42° C), which is also a derivative of acrolein. This unsaturated aldehyde is highly hygroscopic and crystallizes in the form of its hydrate. Its cyclic structure is confirmed by the absence of a reaction with ferric chloride, as well as by analytical and spectroscopic data. The authors also studied synthetic methods for the preparation of 1,2-dimethoxy ethylene (X) which was obtained by the reactions:

 $B_1CH_3CH (OCH_3)_2 \xrightarrow{CH_3OH + N_8} CH_3OCH_2CH (OCH_3)_2$ $H^+ \downarrow (CH_3CO)_2 O$ $CH_3OCH = CHOCH_3 \leftarrow CH_3OCH_2CH OCH_3$ $(X) \qquad (IX) \qquad OCH_3$

Card 3/5

27495 \$/062/61/000/009/011/014 B117/B101

Synthesis and transformations ...

Starting from the acetal of bromo acetaldehyde the authors prepared 1,1,2-trimethoxy ethane (VIII) (b.p. 125°-126°C (741 mm Hg), n25 1.3930) which on boiling with acetic anhydride in the presence of a trace of p-toluenesulfonic acid is transformed to the acylal (IX) (b.p. 64°-65°C (14 mm Hg), n20 1.4055). On treating (VIII) with acetic anhydride and a drop of concentrated sulfuric acid, letting the mixture stand for 14 hr, and then treating it with ice water, with subsequent extraction and distillation, 1,2-dimethoxy-1-acetoxy ethane (IX) was obtained also. The reaction mixture obtained by boiling (IX) for 2 hr at 162°-165°C was slowly distilled from a Favorskiy flask. A mixture boiling at 71°-145°C was obtained. The reaction product was washed with concentrated sodium carbonate solution, dried over potassium carbonate and fractionally distilled, yielding a mixture of isomers in the ratio 3.5: 1 (1,2-dimethoxy ethylene (X), b.p. 97°-99°C, n20 1.4184 and an isomer b.p. 103°C, n20°5 1.4204). The yield of 1,2-dimethoxy ethylene did not exceed 20%. The study of these isomers is being continued. There are 14 references: 3 Soviet and 11 non-Soviet. The four most recent references to English-Card 4/5

s/062/61/000/010/013/018 B106/B101

15.8113

Shostakovskiy, M. F., Khomutov, A. M., and Khomutova, N. M.

AUTHORS: TITLE:

Reaction of polyvinyl alcohol with polymethacrylic acid

PERIODICAL:

Akademiya nauk SSSR. Izvostiya. Otdeleniye khimicheskikh

nauk, no. 10, 1961, 1890 - 1891

TEXT: The activity of the hydroxyl groups of polyvinyl alcohol in the reaction with polymers containing functional groups with mobile hydrogen has hitherto not been studied. In this connection, the authors investigated the reaction between aqueous solutions of polyvinyl alcohol and polymethacrylic acid at room temperature without using a catalyst. The polyvinyl alcohol contained 1.5% of acetate groups, and had a specific weight of 1.259. Data of methacrylic acid: boiling point 160°C, 1.4313, d_4^{20} 1.0153, acid number 650 mg of KOH. Methacrylic acid was polymerized in the presence of 0.2% of benzoyl peroxide, the polymer reprecipitated from its methanolic solution with benzene several times, until no double bond could be proved any longer. Then, polymethacrylic Card 1/4

25276

meaction of polyvinyl alcohol...

S/062/61/000/010/013/016 B106/B101

acid was dried up to a constant weight and finally analyzed. It contained 99.5% of carboxyl groups. In order to synthesize the polyester, 10% aqueous solutions of polyvinyl alcohol and polymethacrylic acid were mixed at room temperature. The polyester precipitate deposited after some minutes was washed with water up to a neutral reaction and then dried up to a constant weight. The content of unused polymethacrylic acid in the filtrate was determined titrimetrically. To analyze the polyester, it was saponified with lye, and the content of carboxyl groups was determined titrimetrically. Then, the amount of polymethacrylic acid entering the composition of the polyester was calculated. The results are given in a table. The reaction in question was conducted at equimolecular ratios of the initial substances (referred to one link) or with an excess of one of the two reactants. In all cases, esterification was almost quantitative. The reaction can be observed well, since both polyvinyl alcohol and polymethacrylic acid are readily soluble in water, whereas the reaction product is not water-soluble and precipitates from the aqueous solution. The reaction follows the

Card 2/4

2: 276 3/062/61/000/010/013/018 B106/B101

Reaction of polyvinyl alcohol...

The resulting new polyesters belong to the so-called cross-linked high-molecular compounds. They are insoluble in water and organic solvents, and carbonize on heating without melting. In order to clarify the structure of the polyesters, they were subjected to alkaline hydrolysis. In aqueous sodium hydroxide, hydrolysis is complete and yields polyvinyl alcohol and the sodium salt of polymethacrylic acid. An analogous expolymethacrylic acid. The solution of the sodium salt of polymethacrylic acid.

periment with an aqueous solution of polyvinyl alcohol was carried out in order to study the course of reaction between polyvinyl alcohol and monomeric methacrylic acid. In this case, the reaction is considerably monomeric methacrylic acid. In this case, the reaction is considerably slower. It was found that the esterification of polymethacrylic acid with polyvinyl alcohol in aqueous solution is almost quantitative.

Abstracter's note: Essentially complete translation. There are table and 3 Soviet references.

Card 3/4

5/062/61/000/010/013/018 B106/B101

ASSOCIATION: Institut organicheskoy khimii im. N. D. Zelinskogo Akademii

nauk SSSR (Institute of Organic Chemistry imeni N. D. Zelinskiy of the Academy of Sciences USSR)

SUBMITTED: April 3, 1961

Reaction of polyvinyl alcohol ...

Исходные вещества	Моляр- нос соотно- шение	Buxog.	Содержин полимета инсл	криловой оты (у)	Количество полимет- акриловой кислоты, (5) не вошед- шей в реак- шкю. %
Поливиниловый спирт © Полиметакриловая кислота Поливиниловый спирт © Полиметакриловая кислота Поливиниловый спирт © Полиметакриловая кислота Полиметакриловая кислота	1 1 0,5 1 1 0,5	99,0 91,6 94,5	61,6 61,6 61,6	59,0 58,6 59,4	1,07 41,6 0,97

Legend to the Table: (1) initial substances; (2) molar ratio; (3) yield, % of theoretical value; (4) content of polymethacrylic acid links; (a) theoretical value, %; (b) practical value, %; (5) amount of polymethacrylic acid which did not react, %; (6) polyvinyl alcohol; (7) polymethacrylic acid.

Card 4/4

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; DENISENKO, V.P.; GORBAN', A.K.

Synthesis of hexamethylenediamine biquaternary ammonium salts. Izv.AN SSSR.Otd.khim.nauk no.10:1907-1908 0 '61. (MIRA 14:10)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Ammonium compounds) (Hexanediamine)

GASTILOVICH, Ye.A.; SHIGORIN, D.N.; GRACHEVA, Ye.P.; CHEKULAYEVA, I.A.; SHOSTAKOVSKIY, M.F.

Investigating the nature of the complexes and derivatives of acetylene by the method of infrared absorption spectra. Opt.i spektr. 10 no.5:595-599 My '61. (MIRA 14:8) (Acetylene—Spectra)

SHOSTAKOVSKIY, M.F.; BOGDANOVA, A.V.; VOLKOV, A.N.

Vinyl compounds in diene synthesis. Report No.8: Diene synthesis of vinyl ethers and thio ethers with anthracene. Izv.AN SSSR.Otd.-khim.nauk no.11:2072-2074 N '61. (MIRA 14:11)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Ethers) (Sulfides) (Anthracene)

29522 S/062/61/000/011/009/012 B103/B147

15-8113

AUTHORS:

Shostakovskiy, M. F., Khomutov, A. M., Chekulayeva, I. A., and

的人,我们就是这种的人,我们就是我们的人,我们就是这个人,我们就是这个人,我们就是这个人,我们就是这个人,我们就是我们的人,我们就是这种的人,我们就会是这种人的

Khomutova, N. M.

TITLE: Synthesis and polymerization of diallyl tartrate

PERIODICAL: Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh

nauk, no. 11, 1961, 2075 - 2077

TEXT: Synthesis and polymerization of diallyl tartrate (DAT) were studied. This was done to clarify the effect of its structure on the course of polymerization as well as the polymer properties, in the case where DAT contains further functional groups (OH). The synthesis was effected by esterification of tartaric acid withoallyl alcohol in the presence of hydroquinone and sulfuric acid at 70 C in benzene solution. In order to establish the polymerization conditions of DAT, different quantities of the following initiators were used: (a) benzoyl peroxide (Bz₂O₂), and (b) azoisobutyric acid dinitrile, the temperature (60, 95, and 125 C) as well as the relation time (5 - 21 hr) being varied. It has been found that

Card 1/1/

29522 · \$/062/61/000/011/009/012 B103/B147

Synthesis and polymerization...

either linear (I) or tridimensional (II) polymers are formed in different quantitative proportions depending on the reaction conditions:

Card 2/# /

29522 5/662 (61/600 (611/600/612 2106/2017

Synthesis and polymerization

Table 'shows the effect of br₂₀₁ as purpletor on the yelymarization of BAT. The polymers have a predominantly cross-linked structure in the presence of 4.5 - 6% of Bz₂O₂. Increase of temperature and heating time have the following effects with 6% of Bz₂O₂. DAT is practically not polymerized within 5 hr at 60°C, whereas a solid and insoluble polymer is formed at 95°C (yield 57%). Such a polymer forms at 60°C only after heating for 21 hr. Polymerization with 6% of Bz₂O₂ at 25°C for 18 hr gave the best results 98% of a solid transparent polymer which cannot be charged by static electricity. Its heat resistance is 294°C. (b) is inferior to Bz₂O₂ as initiator. At 95°C its use yields only 5% of viscous polymer within 18 hr. DAT is less active in polymerization than fumaric and maleic esters. This might be due to the OH groups contained in DAT. There are 2 tables and 6 references. 4 Soviet and 2 non-Soviet. The references to English-language publications read as follows. Tsunao Araki. Hiroko Jida. Repts Govt. Chem. Ind. Research Inst. Tokyo. 47. 95 (1952); Chem. Abstro., 47. 10889 (1953).

Card 3/44

29522 S/062/61/000/011/009/012 B103/B147

Synthesis and polymerization. .

Telinskiy of the Academy of Sciences USSR)

SUBMITTAD: June 12, 1961

Table : Polymerization of diallyl tartrate with different quantities of Bz_2C_2 at 95°C within 5 hr Legend (1) Bz_2C_2 content, β ; (2) fractionated quantity of polymer g; (3) polymer yield, β ; (4) total: (5) tridimensional; (6) linear; (7) quantity of compolymerized monomer β ; (8) content of tartaric and in the tridimensional polymer, β ; (9) real; (10) theoretical

Card 4/1 /

SHOSTAKOVSKIY, M.F.; BOGDANOVA, A.V.; USHAKOVA, T.M.

Vinyl compounds in diene synthesis. Report No.9: Some properties of diene synthesis adducts with vinyl sulfides. Izv. AN SSSR (MIRA 14:11) Otd.khim.nauk no.12:2217-2222 D 61.

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR. (Vinyl compounds)

"APPROVED FOR RELEASE: 08/09/2001

CIA-RDP86-00513R001549920002-9

30466

5/062/61/000/012/007/012 B117/B147

15.8070

Shostakovskiy, M. F., Khomutov, A. M., and Sidel'kovskaya,

Copolymerization of vinyl pyrrolidone with methyl methacrylate AUTHORS: F. P.

and acrylonitrile

Otdeleniye khimicheskikh TITLE: Akademiya nauk SSSR. Izvestiya.

nauk, no. 12, 1961, 2222 - 2225 PERIODICAL:

TEXT: The copolymerization of N-vinyl pyrrolidone with methyl methacrylate and acrylonitrile in various molar ratios up to radical conversion was and adrytonithing in various motar ratios up to radical conversion was examined. Polymerization occurred within 100 hr in the presence of dinitrile of azoisobutyric acid (0.2%) at 60 ± 1°C. During copolymerization of the above monomers, copolymers were formed in which the number of vinyl pyrrolidone groups increased with an increase in concentration of the vinyl pyrrolidone in the reaction medium while the yields slightly decreased The relative activity of radicals of the examined monomers was studied on the copolymerization with lesser degree of conversion. was studied on the copolymerization with leaser degree of constants \mathbf{r}_1 and \mathbf{r}_2 the evaluation of this activity, the copolymerization constants \mathbf{r}_1 and \mathbf{r}_2

Card 1/3

30166

s/c62/61/000/012/007/012 B117/B147

Copolymerization of vinyl pyrrolidone...

were determined with an accuracy of \pm 0.02 using the integral equation of Mayo and Lewis (Ref. 3, see below). A comparison of the relative activities showed that methyl methacrylate was more active with respect to vinyl pyrrolidone radicals. To clarify the effect of vinyl pyrrolidone groups on the solubility of copolymers with acrylonitrile groups, the solubility of the copolymers in several organic solvents was examined at rcom temperature and during heating. It was found that copolymers of vinyl pyrrolidone and methyl methacrylate were soluble in acetone, ethanol, butanol, benzene, dioxane, chloroform, ethyl cellosolve, ethyl acetate, and butyl acetate. Copolymers of vinyl pyrrolidone and acrylonitrile were not soluble in the above-mentioned compounds. They dissolve in pyrrolidone, vinyl pyrrolidone, butyl pyrrolidone, butyrolactone. β (N-pyrrolidonyl) ethyl formiate, β (N-pyrrolidonyl)-ethyl acetate. the above mentioned organic compounds, the homopolymer of acrylonitrile is insoluble. There are 2 figures, 5 tables, and 5 references: 2 Soviet and 3 non-Soviet The three references to English-language publications read as follows: Ref. 1: U. S. Pat. 2667473 (1954); U. S. Pat. 2676949 (1954); U. S. Pat. 2497705 (1950); U. S. Pat. 27:3573 (1955); U. S. Pat. 2739588 (1956); R. M. Rike , D. L. Baily, J. Polymer Sci. 22, no. 100, 55

Card 2/3

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; KALABINA, A.V.; KOMAROV, N.V.

Synthesis and transformations of vinyl aryl ethers. Report No.1: Synthesis and properties of vinyl ether of p-sec-propylphenol. Izv. Fiz.-khim. nauch.-issl. inst. Irk. un. 5 no.1:215-224 '61. (MIRA 16:8)

(Ethers) (Phenol)

SHOSTAKOVSKIY, M.F.; SIDEL'KOVSKAYA, F.P.

Medicinal preparations based on polymers. Med. prom. 15 no.3:6-13 Mr '61. (MIRA 14:5)

1. Institut organicheskoy khimii imeni N.D.Zelinskogo AN SSSR. (POLYMERS) (DRUGS)

SOSTAKOVSKI, M. F. [Shostakovskiy, M. F.]; SIDELKOVSKAIA, F. P. [Sidel'kovskaya, F. P.]

Drugs on the polymer basis. Analele chimie 16 no.4:21-30 C-D 161.

(Drugs) (Polymers and polymerization)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; GRACHEVA, Ye.P.; KUL'BOVSKAYA, N.K.

Methods for synthesizing and properties of substituted vinyl ethers and substituted vinyl sulfides. Usp. khim. 30 no. 4:493-516 Ap 161. (MIRA 14:4)

l. Institut organicheskoy khimii imeni N.D. Zelinskogo AN SSSR. (Sulfides) (Ethers)

THE REPORT OF THE PROPERTY OF

89522

15.8116

8/079/61/031/002/018/019 B1 18/B208

AUTHORS:

Shostakovskiy, M. F. and Vlasov, V. M.

TITLE:

Synthesis of trialkyl vinyl hydroxy silanes from acetylene and

Zhurnalobshchey khimii, v. 31, no. 2, 1961, 683 PERIODICAL:

TEXT: Vinyl ethers containing silicon recently raised the interest of chemists being applicable as monomers to the synthesis of high-molecular compounds. The authors were the first to synthesize trialkyl vinyl hydroxy silanes, and to describe their properties (Ref. 1). Other authors showed in Ref. 2 that trialkyl vinyl hydroxy silanes may be synthesized by another method. As the attempts of the afore-mentioned research workers of synthesizing these compounds by the method of direct vinylation according to the reaction of Favorskiy-Shostakovskiy were unsuccessful, the incorrect conclusion was drawn that this reaction be not applicable to the synthesis of trialkyl vinyl hydroxy silanes (Ref. 3). Thorough investigations of the conditions to be observed in direct vinylation carried out by the authors disclosed that the main hindrance in this reaction, under the conditions of Card 1/2

89522 s/079/61/031/002/018/019 B118/B208

Synthesis of trialkyl ...

synthesis, was the high tendency of silanols toward conversion to disiloxane To suppress this undesirable conversion, they suggested the application of excess acetylene. The synthesis of trialkyl vinyl hydroxy silanes was rendered possible according to equation

 $(R)_3$ SiOH + nHC = CH \longrightarrow $(R)_3$ Si - O - CH = CH₂ + n - 1HC = CH.

[Abstracter's Note: This is a complete translation of the original paper.]

ASSOCIATION: Institut organicheskoy khimii Akademii nauk SSSR (Institute of

Organic Chemistry of the Academy of Sciences USSR)

SUBMITTED: September 18, 1960

Card 2/2

2209, 1287, 153 5.3630

21082 s/079/61/031/003/002/013 B118/B207

AUTHORS:

Shostakovskiy, M. F., Shmonina, L. I., and Guseynov. I. I.

TITLE:

Synthesis of 1-alkoxy-2-oxy-dialkoxy- and 1-alkoxy-2-oxy-bisdialkyl aminophosphine-3-chlorobutadiene-1,3

PERIODICAL:

Zhurnal obshchey khimii, v. 31, no. 3, 1961, 734-736

TEXT: The new class of 1,2,3-substituted butadienes (I) previously described by the authors (Ref. 1: ZhOKh, 30, 2836 (1960)), which was obtained by reacting vinyl-ethinyl alkyl ether with PCl5, contains active chlorine atoms in

its group of oxychloro phosphines:

 $ROCH = CH - C = CH + 2PCl_5 \longrightarrow [ROCHCl - CHPCl_4 - C = CH \cdot PCl_5] \xrightarrow{SO_2}$

→ ROCH=CPOC1, -- CC1= CH₂.

The reaction of butadienes (I) with compounds containing a mobile hydrogen atom (alcohols, amines, mercaptanes) may play a Card 1/3

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CIA-RDP86-00513R001549920002-9"

21082 S/079/61/031/003/002/013 B118/B207

Synthesis of ...

considerable role in the synthesis of valuable polymers, if the halogen atoms of the phosphine group are substituted by different radicals. In the reaction of compounds (I) with alcohols, e. g., the chlorine atoms of the phosphine group are easily substituted by alkoxy groups under the formation of 1-alkoxy-2-oxy-dialkoxy phosphine-3-chlorobutadiene-1,3 (II): ROCH= CPOC1₂-CC1= CH₂ + 2R'OH - ROCH= CPO(OR')₂-CC1= CH₂ (II). This reaction proceeds easily already at 30-35°C in anhydrous benzene, in the presence of pyridine. The yellowish liquids obtained have a weak odor and are more stable than the initial butadienes (I) and may be preserved in sealed ampoules. Compounds (I) react with dialkyl amines in the same way. In this case, the reaction proceeds in anhydrous petroleum ether at -3°C, in the presence of excess dialkyl amine. In analogy to the compounds (I) and (II), the resulting 1-alkoxy-2-oxy-bis-dialkyl aminophosphine-3-chlorobutadienes-1,3 are assigned the structure ROCH=CPO(NR'2)2-CC1-CH2 (III). They are dense, yellowish liquids, which become darker when exposed to air. firm the previously assumed structure of 1.2,3-substituted butadienes (I), their Raman spectra were examined. The oscillation frequency of the C-Cl bond in the grouping - CCl= CH2 was found to conform to formula (I),

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Synthesis of ...

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and the compounds studied revealed a rotational isomerism at the single C-C and C-P bonds, as well as a geometrical isomerism at the C-C double bond which follows the C-P bond. The infrared spectra of some compounds (II) showed an absorption band 1408-1400 cm-1 which corresponds to a vinyl group (= CH₂) in final position, which fact is also in agreement with the structure of 1,2,3-substituted butadienes of these compounds and spectroscopic data presented in Ref. 1. There are 4 Soviet-bloc references.

ASSOCIATION:

Institut organicheskoy khimii Akademii nauk SSSR (Institute of Organic Chemistry of the Academy of Sciences USSR)

SUBMITTED:

April 21, 1960

Card 3/3

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

SHOSTAKOVSKIY, M.F.; RABINOVICH, M.S.; LEVITOV, M.M.; VERKHOVTSEVA, T.P.; PREOBRAZHENSKAYA, Ye.V.; KULIKOVA, G.N.; KALINOVSKIY, O.A.

Synthesis of the precursous and fragments of antibiotics. Part 4: Thioglycolic acid derivatives. Zhur.ob.khim. 31 no.5:1453-1458 My ¹61. (MIRA 14:5)

1. Vsesovuznyy nauchno-issledovateliskiy institut antibiotikov. (Acetic acid) (Antibiotics)

"APPROVED FOR RELEASE: 08/09/2001 CIA-RDP86-00513R001549920002-9

KOSTROMINA, O.Ye.; SHOSTAKOVSKIY, M.F.; VASIL'YEVA, N.Ye.

Synthesis of the precursors and fragments of antibiotics. Part 5:
Synthesis of di- and trihaloderivatives of cyclohexenecarboxylic acid. Zhur.ob.khim. 31 no.5:1458-1462 My '61. (MIRA 14:5)

1. Vsesoyuznyy nauchno-issledovatel'skiy institut antibiotikov. (Cyclohexenecarboxylic acid)

(Antibiotics)

SHOSTAKOVSKIY, M.F.; VORONKINA, T.M.; SIDEL'KOVSKAYA, F.P.

Synthesis of the precursors and fragments of antibiotics. Part 6:
Derviatives of lactam-containing mercaptocetic acid. Zhur.ob.khim.
31 no.5:1463-1465 My 161. (MIRA 14:5)

l. Vsesoyuznyy nauchno-issledovateliskiy institut antibiotikov.
(Acetic acid) (Antibiotics)

SHOSTAKOVSKIY, M.F.; PRILEZHAYEVA, Ye.F.; AZOVSKAYA, V.A.; YANSONS, I.;
YANSONS, S.

Diene syntheses with bivinyl sulfur-containing dienophiles.
Zhursob.khim. 31 no.6:2079-2080 Je *61. (MIRA 14:6)

1. Institut organicheskoy khimii AN SSSR. (Dienophiles) (Butadiene)